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Dated: October 6, 2005	Signature:	()

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

App. No.

10/516,864

Confirmation No. 8549

Applicant

Hsiao, et al.

Filed

June 27, 2003

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Examiner

Not assigned

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Docket No.

32144183-000004

Customer No.:

51738

Entitled

Plasma or Serum Marker and Process for Detection of Cancer

Commissioner for Patents

P.O. Box 1450

Alexandria, VA 22313-1450

DECLARATION OF MICHAEL DAI UNDER 37 CFR §1.47

- I, Michael Dai, Declare as follows:
- 1. I am at least 18 years of age and am competent in all respects to make the following statements.
- 2. I represent Hong Kong University of Science and Technology for US Patent Application No. 10/516,864.
- 3. Exhibit A is a statement regarding Lack of Signature for inventor Cesar Wong who refused to sign the Declaration of Inventorship for the purposes of the designation of the United States of America and also refused to sign the PCT Power of Attorney.
- 4. Exhibit B is an e-mail dated August 26, 2005 to Cesar Wong attached thereto a copy of the Declaration and a fullcopy of the current application, requesting the signing of the declaration form by Cesar Wong.

5. Exhibit C is a return read receipt e-mail documenting that the aforesaid email has been displayed on the recipient's computer.

6. Exhibit D is a Delivery Services Request Form showing that on August 30, 2005 around

9:00pm, our firm's representative visited Cesar Wong's last known address to deliver a

confirmation copy of our August 26 e-mail together with a copy of the declaration and a full

copy of the patent application to Cesar Wong. However, we were informed by the lady who

answer the door and by the building security guard that Cesar Wong no longer lives there.

7. Exhibit E1 is a copy of the DHL shipment form showing that on September 2, 2005, we

arranged for DHL Domestic Shipment to deliver the confirmation copy of our August 26

email together with copies of the declaration and the patent application to Cesar Wong's

working address. Exhibit E2 is a copy of the DHL online report showing that the delivery

was completed on September 2, 2005. Exhibit E3 is the fax copy of the DHL Notification

Form confirming the same.

8. I further declare that all statements made herein of my own knowledge are true and that all

statements made on information and belief are believed to be true; and further that these

statements were made with the knowledge that willful false statements and the like so made

are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United

States Code and that such willful false statements may jeopardize the validity of any

application for which it is used.

Dated: September 7, 2005

Michael Dai,

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Attorneys For Applicants

$\underline{\text{Exhibit } A}$

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Statement Regarding Lack of Signature
PCT Rule 4.15(b) states:
Where two or more applicants file an international application which designates a State whose national law requires that national applications be filed by the inventor and where an applicant for that designated State who is an inventor refused to sign the request or could not be found or reached after diligent effort, the request need not be signed by that applicant if it is signed by at least one applicant and a statement is furnished explaining, to the satisfaction of the receiving Office, the lack of the signature concerned.
Applicant/inventor, Sze-Chuen Cesar Wong, after diligent efforts, has refused to sign the Declaration of Inventorship (Rules 4.17(iv) and 51bis.1(a)(iv)) for the purposes of the designation of the United States of America. In addition, Applicant/inventor, Sze-Chuen Cesar Wong, after diligent efforts, has refused to sign the PCT Power of Attorney.

Dai, Michael

From: Cheung, Peggy

Sent: Friday, August 26, 2005 5:49 PM

To: 'cesar01@netvigator.com'

Cc: 'ttalice@ust.hk'; 'rockylaw@ust.hk'; Dai, Michael; Mok, Chindy

Subject: HKUST - US Patent Application "Plasma or Serum Marker and Process

for Detection of Cancer" (Our Ref: 32144183-000004)

Importance: High

Attachments: .4DECLARATION-POA COMBINED.pdf; PCT specification + amended

claims.PDF

The Hong Kong University of Science & Technology US (PCT) Application No. 10/516,864

Title: Plasma Serum Marker and Process for Detection of Cancer

Dear Mr. Wong,

We represent The Hong Kong University of Science & Technology ("HKUST") in the prosecution of the above referenced patent application, which you are named as an inventor.

The application has entered the national stage in the US. According to the US Patent Law and regulations, all named inventors need to sign a declaration form declaring that he/she is the inventor of the claimed inventions in the patent application. A full copy of the filed patent application is attached hereto for your review and reference. For your ease of handling, we also attach hereto a filled-up declaration form for your review and execution. Please print out a copy of the declaration form and sign at the space provided therein. Please send the declaration form bearing your original signature back to us as soon as possible.





.4DECLARATION-P PCT specification + A COMBINED.pdf... amended cl...

A confirmation copy of this email together with the attachments will be forwarded to you by mail to the address shown in the declaration form. If the address is incorrect, please let us know by return e-mail. Thank you.

If for some reasons you do not want to sign the declaration, or you believe there are conditions that prevent you from signing the declaration, please let us know. If you desire to have further discussion with us regarding the patent application/or and the declaration, please feel free to contact us at the numbers and addresses provided below.

We look forward to hearing from you.

Thanks & regards, Peggy / Michael

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My residence, mailing address, I believe I am the original and fir					hich a pat	ent is sought	on the inver	ntion entitled:
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I hereby claim foreign priority benefits under 35 U.S.C. 119(a)-(d) or (f), or 365(b) of any foreign application(s) for patent, inventor's or plant breeder's rights certificate(s), or 365(a) of any PCT international application which designated at least one country other than the United States of America, listed below and have also identified below, by checking the box, any foreign application for patent, inventor' or plant breeder's rights certificate(s), or any PCT international application having a filing date before that of the application on which priority is claimed.								
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Address							
City		State	ZIP				
Country	Telephone 214/97	78-3000	Fax 214/978-3099				
I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under 18 U.S.C. 1001 and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.							
NAME OF SOLE OR FIRST INVENTOR:	A petiti	ion has been filed for this unsigned in	ventor				
Given Name Sze-Chu (first and middle [if any])	1						
Inventor's Signature			Date				
Happy Valley Residence: City	State	Hong Kong Country	CN Citizenship				
Flat C, Floor 9, King's Court	t, 14-16 Village R	₹oad					
city Happy Valley	State	ZIP	Country Hong Kong				
NAME OF SECOND INVENTOR:	A petition has bee	en filed for this unsigned inventor					
Given Name (first and middle [if any])		Family Name or Surname					
Inventor's Signature			Date				
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Additional inventors are being named of	on the supplemental Add ⁱ	itional Inventor(s) sheet(s) PTO/SB/0					

PLASMA OR SERUM MARKER AND PROCESS

FOR DETECTION OF CANCER

CROSS-REFERENCE TO RELATED APPLICATION

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This non-provisional International Patent Application claims priority from U.S. Provisional Application Serial No. 60/392,191, filed on June 28, 2002, and entitled "Plasma or Serum Marker and Process for Detection of Cancer", which is commonly owned with the present application and incorporated herein by reference for all purposes.

FIELD OF THE INVENTION

The present invention relates to a PCR based process in detection of blood plasma or serum marker for diagnosis, early detection, monitoring and population screening for cancer and, more particularly, detection of β -catenin RNA and DNA in blood plasma or serum for colorectal cancer.

BACKGROUND OF THE INVENTION

Colorectal cancer (CRC) is one of the most common malignancies worldwide. The number of new cases of CRC has been increasing rapidly since 1975. More than 70% of CRC cases develop from sporadic adenomas or adenomatous polyps. Early detection and surgical removal of polyps is believed to be the most effective way to prevent benign polyps from developing into malignant tumors and thereby reducing mortality caused by CRC.

Traditional screening methods for colorectal cancer include sigmoidoscopy, fecal occult blood testing, colonscopy and double contrast barium enema. However, these traditional methods suffer from limitations and are invasive, high cost, of low predictive value or result in low detection rates. For example, WO0142504, the teachings of which are incorporated herein by reference, discloses a multi-reaction process for detection of extracellular tumor associated nucleic acid in blood plasma or serum. Further advances are desirable.

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β-catenin protein was initially identified through its interaction with cadherins. Recent evidence shows that it acts as a transcriptional factor and plays a key role in the Wnt-signaling pathway (Willert & Nusse, 1998). It has been demonstrated that accumulation of cytoplasmic and nuclear β-catenin signaling is tightly associated with the genesis of a wide variety of tumors. (Morin, 1999).

It has been discovered that using immunohistochemical staining that levels of nuclear β -catenin are highly correlated with the purported sequential stages in colorectal carcinogenesis with positive staining observed in 0% of normal tissues, 8% of polyps, 92% of adenomas and 100% of carcinomas. It has been further discovered that the nuclear β -catenin signal appears to clearly differentiate the polyps (non-adenomatous polyps) from adenomas (adenomatous polyps). This would be a useful marker for clinical diagnosis, or early detection of CRC, with the adenoma being considered as endpoint for risk factor. However, this diagnostic method based on the evaluation of nuclear β -catenin requires colonscopic procedure, then surgical removal of the suspected tissues.

Accordingly, there is a need for an effective, less invasive, more accurate test for early detection of cancer. The present invention meets this need.

SUMMARY OF THE INVENTION

The present invention provides a PCR (Polymerase Chain Reaction) based method or process in the detection of serum or plasma marker RNA and DNA related to β -catenin providing an effective, less evasive and more accurate test for the diagnosis, early detection, monitoring, and population screening of colorectal and other cancer types. It will be appreciated that this method of detection of β -catenin RNA and DNA in blood serum can be applied to other plasma and serum RNA and DNA encoded for β -catenin associated proteins. In one embodiment, the RNA or DNA is derived from genes encoded beta-catenin, alpha-catenin, E-catherin and other beta catenin associated proteins.

The process of the present invention comprises detecting blood serum or plasma RNA or/and DNA from a human or animal as a tool in the diagnosis, early detection, monitoring, treatment and population screening of neoplastic diseases at various progression and clinical stages. One advantage of the present invention is the

non-invasive nature of the method, and a second advantage is improved accessibility of sample collections and sensitivity

Details of multiple embodiments of the invention are set forth below. These embodiments are for illustrative purposes only and the principles of the invention can be implemented in other embodiments. Other features and advantages of this invention will become apparent from the following description and examples.

BRIEF DESCRIPTION OF THE DRAWINGS

For a more complete understanding, reference is now made to the following detailed description taken in conjunction with the accompanying drawings. It is emphasized that some components may not be illustrated for clarity of discussion. Reference is now made to the following descriptions taken in conjunction with the accompanying drawings, in which:

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- FIG. 1a, FIG. 1b and FIG. 1c illustrate detection of β-catenin RNA from plasma of colorectal carcinoma patients using RT-PCR.
 - FIG. 2a and FIG. 2b illustrate detection of blood β -catenin RNA from patients for colorectal adenoma using RT-PCR.
- FIG. 2c illustrate detection of blood β-actin RNA from patients for colorectal adenoma using RT-PCR.
- FIG. 3a, FIG. 3b, FIG. 3c, FIG. 3d, and FIG. 3e illustrate detection of serum β-catenin DNA from patients with adenomas or carcinomas and normal controls.

DETAILED DESCRIPTION OF THE INVENTION

The search for sensitive and specific biomarkers for early detection of colorectal cancer has been discovered in the present invention. The advanced understanding of the molecular mechanism underlying the carcinogenesis of colorectal cancer has helped to identify a few oncogenes and tumor suppressors as potential clinical biomarkers of colorectal cancer development and early detection. These include k-ras, APC, p53, MCC, DCC genes. However, none of the candidate

condensed or omitted altogether inasmuch as detail discussions of these features are not considered necessary to obtain a complete understanding of the disclosure, and are considered to be within the understanding of persons of ordinary skill in the relevant field of art.

Example 1

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*β-catenin RNA was detected in all plasma samples of patients with colorectal carcinoma.

To detect the presence of plasma β-catenin, RT-PCR (reverse transcription-polymerase chain reactions) were preformed on two blood samples from patients with carcinoma using Primer #1 that would yield a 224 bp of exon 3 region of the gene. An RNA sample extracted from carcinoma tumor expressing high level of β-catenin was included as positive control. Results showed that two plasma and the positive control RNA samples yielded a 224 bp band in the presence, but not in the absence of, reverse transcriptase (RT) in the reaction (FIG. 1a). RT-PCR analysis was preformed on the other 10 plasma RNA samples using the intron spanning primers (Primer#2, Table 2).

Data showed that a 250 bp fragment was clearly detected in all 10 patient plasma samples (FIG. 1a, lanes 1-10), suggesting the presence of β-catenin RNA in the circulating blood of carcinoma patients. The data also showed that the reaction is RT-dependent (FIG. 1b, lane 12). A genomic DNA sample was included as a positive control for PCR reaction and a 450 bp band appeared as expected (FIG. 1b, lane 11).

To prove that the 250 bp band was derived from the RNA, instead of DNA templates in the plasma, tests were performed on the three remaining plasma RNA samples without prior treatment with DNase I.

Two PCR products, a 250 bp band amplified from RNA and a 450 bp band amplified from the DNA contaminating plasma RNA extract, appeared on the gel. All three samples yielded both 250 and 450 bp bands in the presence of RT (FIG. 1b, lanes13-15), and a single 450 bp band was observed from a RNase treated DNA sample in the absence of RT (FIG. 1b, lane 1).

Fifteen patients were tested with carcinoma using three slight different experimental settings described above, and the data showed that 15 in 15 patients were clearly positive for plasma β -catenin.

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Example 2

*Plasma RNA was present at high rates in patients with adenomas, but not in healthy individuals.

Seventeen plasma samples were screened for β-catenin RNA from individuals with suspected adenomas. Of the 17 plasma samples from individuals with suspected adenomas screened for β-catenin RNA, 11 were plasma positive, indicated by the presence of a 250 bp RT-PCR product; 6 were found negative (FIG. 2a, lanes 1-11; FIG. 2b, lanes 1-6). RT-PCR assays were performed on the 6 negative samples using primers specific for β-actin sequences (Table 2, Primer#3). β-actin RNA was detected in all six plasma samples (FIG. 2c, lanes 1-6), indicating the six plasma RNA extracts were in amplifiable quality. Of the 6 patients with negative β-catenin signals (Table 1, Patients#10, 14, & 16), biopsy later confirmed that three were diagnosed with adenoma, two had granulation tissues, and the other had a dilated lymphatic space (Table 1, Patients#1-3). The percentage of detection among adenoma patients was 79% (11 of 14). Parallel RT-PCR analyses were performed on 10 healthy subjects. Nine of the ten healthy controls showed negative plasma β-catenin signals, but all showed positive β-actin RNA signals (FIG. 2d & FIG 2e, lanes 1-10). Only 1 of them had a rather weak positive signal (FIG. 2d, lane 10).

In summary, the presence of β -catenin was examined in the blood plasma of 32 patients with confirmed carcinoma or adenoma using RT-PCR analysis. Results showed that 100% (15 of 15) of patients with carcinoma, 79% (11 of 14) of patients with adenoma and 10% (1 of 10) healthy volunteers carried β -catenin RNA in their circulating blood. It is worthy to mention that the apparently healthy subject with weak plasma β -catenin RNA had been suffered from long-standing colorectal discomfort, occasionally with fecal blood and diarrhea, although no abnormality or ulceration colitis was detected in an endoscopic examination. Three patients with suspected adenoma at admission were also tested for plasma β -catenin. All three patients who were later confirmed by biopsy to be free of adenoma were negative for plasma signal.

It has been shown that free DNA is present in the circulating blood of patients with disorders and cancers, and this DNA can be detected using PCR assay.

Furthermore, reports have showed that genetic alterations of specific gene sequences can be detected in the serum of cancer patients (Anker P 1997; Hibi K

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Example 3

*Immunochemical staining of nuclear β -catenin signals of the adenoma and carcinoma tissues.

In more than 200 cases examined, 92% of adenomas and 400% of carcinomas, but none of the normal tissues showed elevated nuclear β -catenin. To determine the nuclear β -catenin signals of the adenomas and carcinomas obtained from patients derived from Examples 1 & 2, paraffin-embedded tissue blocks of adenoma and carcinoma of 32 patients were sectioned and examined for nuclear β -catenin. The immunohistochemical staining was scored based on both the intensity and the percentage positive cells. Table 2 showed that nuclear translocation of β -catenin was observed in all tissue specimens.

Example 4

*Quantification of blood β-catenin RNA in healthy individuals and patients with adenoma or carcinoma using real time RT-PCR technology.

The quantitative difference in plasma β -catenin signal between adenoma and carcinoma patients was investigated using real-time reverse transcriptase-PCR (RT-PCR). The results showed that the average copy number of β -catenin mRNA was 30 fold higher in adenoma (n=12; 3 negative; 8 positive: mean, $1.1x10^3$; ranging from $0.69x10^3$ to $1.80x10^3$) and 598 fold higher in carcinoma (n=18; mean, $2.2x10^4$ ranging from $.67x10^4$ to $4.4x10^4$) patients than the normal individuals ((n=14; mean, 36 ranging from 0 to 169). The copy number of β -catenin mRNA in carcinoma patients was 19 fold higher than in adenoma patients. These quantification analysis provide a clear evidence that the plasma β -catenin mRNA are present differentially and can be used as a diagnostic tool to differentiate healthy subject, adenoma and carcinoma patients.

Example 5

*Detection of β-catenin DNA in the serum of patients with colorectal adenoma and carcinoma.

PCR analysis was first performed with serum DNA samples extracted from colorectal carcinoma patients. The results showed that a 359 bp band was observed in

all 15 serum DNA samples (FIG. 3a, lanes 1 to 16). Ten patients were tested with confirmed adenoma ranging from mild to severe dysplasia. Positive band was detected in 9 of 10 patients (FIG. 3b, lanes 1-11). The detection rate was 90%. The only negative case (FIG. 3b, lane 8) was amplifiable as it yielded positive 156 bp band after amplification with RET specific primers (FIG. 3d, lower panel, lane 13). PCR amplification of β -catenin was also performed on 10 healthy volunteer controls. None of the serum samples showed positive signals for β -catenin, while positive signals were clearly detected using RET specific primers (FIG. 3c, lanes 1 to 10; & 1D, lanes 1-11). In addition, a known positive carcinoma serum sample was carried out in parallel and showed typical 359 bp band on the agarose gel (FIG. 3c, lane 11). Lane 12 of FIG. 3c & FIG. 3d are the negative control for PCR reaction.

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The data showed, for the first time, that serum β-catenin DNA is detectable in all patients with colorectal carcinoma and in 9 out of 10 patients with colorectal adenoma, while all 10 healthy individuals were free of serum \(\beta\)-catenin DNA. This result suggests that the presence of β-catenin DNA in the blood is significantly correlated with the existence of cancer at both preneoplastic and malignant stages. which may also suggest that the circulating β-catenin originated from the adenoma or carcinoma tissue of the patients. The ten adenoma patients, the individual (Patient #9, Table 4) negative in serum β-catenin had the smallest adenoma in this example (3.5 mm in diameter, 48 mm³)). Patient with the next smallest size of adenoma (63 mm³) showed PCR amplifiable \(\beta\)-catenin DNA in the blood, suggesting that the sensitivity of the current method would allow us to detect premalignant adenomotous polyps at least as small as 63 mm³. Quantification of the copy number of β-catenin DNA in the samples using real-time PCR analysis is suggested. The findings indicate that measuring the levels of β -catenin DNA in the blood provides a highly sensitive but noninvasive method for early detection of colorectal cancer. This method may be extended to cancers of different tissue origins.

Referring now to the drawings, FIG. 1 collectively shows detection of β -catenin RNA from plasma of colorectal carcinoma patients using RT-PCR. More specifically, FIG. 1a shows RT-PCR amplification of β -catenin using β -catenin exon primers. Lanes 1-4, RT-PCR reactions of blood RNA samples isolated from two carcinoma patients in the presence (Lane 1 & 3) and absence (Lane 2 & 4) of RT

enzyme; Lane 5, mRNA extracted from carcinoma specimen expressing β -catenin as a positive control; Lane 6, a buffer control. M: RNA markers. FIG. 1b shows RT-PCR amplification of β -catenin using β -catenin intron-spanning primers. Lanes 1-10, DNAase-treated plasma RNAs isolated from ten carcinoma patients; Lane 11, genomic DNA as a positive control for PCR reaction; Lane 12, a buffer control. Lanes 13-17, Samples derived from Lanes 8-12 respectively without prior DNAase treatment. FIG. 1c shows Lane 1-3, β -catenin RNA (250 bp) isolated from three patients by RT-PCR with intron-spanning primers without DNAase treatment; lane 4, positive DNA control; lane 5, negative buffer control. M: DNA markers.

FIG. 2 shows detection of blood β-catenin (FIG. 2a & FIG. 2b) & β-actin (FIG. 2c) RNA from patients suspicious for colorectal adenoma (FIG 2a-2c) using RT-PCR. A. Lanes 1-17, plasma RNAs isolated from 17 patients; Lane 18, positive DNA control; Lane 19, negative control. Detection of blood β-catenin (FIG 2d) & β-actin (FIG 2e) RNA from plasma of ten healthy objects (Lanes 1-10). Lane 11, positive DNA control, Lane 12, negative buffer control.

FIG. 3 shows detection of serum β-catenin DNA from patients with adenomas or carcinomas and normal controls. FIG. 3a, FIG. 3b and FIG. 3c show PCR analyses with β-catenin specific primers were performed with serum samples isolated from patients with colorectal carcinoma: FIG 3a, lanes 1-15; with colorectal adenoma: FIG 3b, lanes 1-10; from healthy individuals: FIG 3c, lanes 1-10. FIG 3d: PCR reactions with RET specific primers were performed with serum samples with negative β-catenin signal. Lanes 1-10, same healthy individual serum samples shown in FIG 3c; FIG 3d, lane 13: the same serum sample shown in Panel FIG 3b, lane 8. Positive control genomic DNA isolated from carcinoma tumor: FIG 3a, lane 16; FIG 3b, lane 11; FIG 3c, lane 11; FIG 3d, lane 11. Negative cell free control: FIG 3a, lane 17; FIG 3b, lane 12; FIG 3c, lane 12; FIG 3d, lane 12. M: Hae III λ DNA marker.

Techniques applied:

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Blood samples and RNA extraction

A 6-ml blood sample was collected from each patient by transcutaneous needle into 8-ml Vacutaniners containing EDTA lithium heparin. Blood samples were centrifuged at 4800 rpm for 8 min. Plasma was aliquoted into polypropylene tubes

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and stored at -80°C for later RNA extraction. RNA was extracted from plasma sample using TRIZOL Kit (Life Technologies, USA), then purified with RNeasy column (Qiagen, Germany) according to the manufacturer's manuals. In brief, 2ml of each plasma sample was mixed with 1.6 ml TRIZOL and 0.4 ml chloroform, centrifuged at 12,000 rpm for 15 min at 4°C. The aqueous phase was collected for RNA extraction using the RNeasy column. The isolated RNA was dissolved in 15 µl of DEPC-treated water. The RNA samples were further treated with PCR grade of deoxyribonuclease I (DNase I)(Life Technologies). In the reaction, 1 µl each of 10 x DNase I reaction buffer and DNase I were added into the 15 µl of RNA sample and incubated at room temperature for 15 min followed by inactivation of DNase I by the addition of 1 µl of 15 mM EDTA and heated at 65°C for 5 min, then chilled in ice before RT-PCR reaction.

Primers and RT-PCR reactions of blood RNA samples

The detection of plasma β-catenin was performed using RT-PCR assay with a set of primers including intron sequence spanning between exon 3 and 4 of β-catenin gene (Table 1). For comparison, a separate set of primers sequences within exon 3 of the β-catenin gene was also incorporated in some PCR reactions. The reverse transcription reaction was performed according to the manufacturer's guides (Qiagen, Germany). PCR was carried out using reagents supplied in a GeneAmp DNA Amplification Kit using AmpliTaq Gold as the polymerase (Perkin-Elmer Corp., Foster City, CA). The parameters used in PCR were 40 cycles with initial denaturation at 95°C for 10 min, followed by 94°C for 1 min 15 s, 59 °C (β-catenin) for 1 min 30 s, 72°C for 1 min 30 s, with a final extension step of 72°C for 10 min. PCR products were analyzed by 1.5% agarose gel electrophoresis and ethidium bromide staining. A negative (water) control was included in each RT-PCR assay. All samples with negative results were subjected to RT-PCR assay for β-actin RNA using intron-spanning primers (Table 3) as a control for the amplifiability of plasma-extracted RNA.

DNA extraction

Blood sera were removed from the supernatants of clotted blood samples and were centrifuged at 4800 rpm for 8 minutes, followed by gently aliquoting of serum

into polypropylene tubes and storage at -20°C for later DNA extraction. DNA was isolated from 200 µl serum using QIAamp DNA Mini Kit (Qiagen, Hilden Germany) according to the manufacturer's protocol. The DNA samples,, were eluted with 50 µl of ddH20.

Primers and PCR reactions of blood DNA samples

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The detection of β-catenin was performed using PCR assay with set of primers franking the 2nd and the 3rd introns of β-catenin gene (Table 3). The PCR was carried out using reagents supplied in a GeneAmp DNA Amplification Kit using AmpliTaq Gold as the polymerase (Perkin-Elmer Corp., Foster City, CA). The parameters used in PCR were 40 cycles with initial denaturation at 95°C for 10 min, followed by 94°C for 1 min and 15 s, 57°C (β-catenin) and 69°C (RET) for 1 min 30 s, 72°C for 1 min 30 s, with a final extension step of 72°C for 10 min. PCR products were analyzed by 1.5% agarose gel electrophoresis and ethidium bromide staining. PCR products were confirmed by direct DNA sequencing. A negative (water) control was included in each PCR assay. All samples with negative results were subjected to PCR assay for RET gene as a control for the amplifiable quality of the serum DNA samples. The RET gene sequence which encodes receptor tyrosine kinase, is normally present in circulating blood of healthy individuals (Matisa-Guiu 1998).

Immunohistochemical staining and evaluation

Monoclonal antibody to β-catenin (C19220) was purchased from Transduction Laboratories (U.S.A.). The antibody was produced against the C-terminal of a mouse β-catenin protein (a.a. 571-581), and is reactive to β-catenin of human, rat and mouse species. Tissue sections with 4 μm thickness were placed on silane-coated (Sigma Chemicals, St. Louis, MO) glass slides, air dried overnight and rehydrated with xylene and graded alcohol. Antigen retrieval and immunochemical staining was performed in the Ventana-ES automated immunostainer (Ventana, Tucson, Az) as described. The sections were counterstained with Harris haematoxylin and mounted with permount after dehydration in graded alcohol. The negative control was done by replacing β-catenin antibody with TBS. Positive signals were evaluated in 4 fields under a light microscope at 10x40 magnification, without knowledge of the clinical

Table 1: Sequence of primers used in the PCR reactions.

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	Prim size	er N	lucleotide sequence (5' to 3')	Design	product
5	1	sense: antisense	ATTTGATGGAGTTGGACATGG : AGCTACTTGTTCTTGAGTGAA	Within exon 3 of β-Catenin ge	ne 224 bp
	2	sense: antisense	TGATTTGATGGAGTTGGACAT : CATTGCATACTGTCCATCAAT	Intron-spanning between exon 3 & 4 of β -Catenin gene	DNA: 450 bp cDNA: 250 bp
10	3	sense: antisense	AAATCGTGCGTGACATTAAGG : ATGATGGAGTTGAAGGTAGTT	Intron-spanning between exon 4 & 5 of β-actin gene	DNA: 324 bp cDNA: 230 bp

Table 2. Correlation of plasma β -catenin RNA in colorectal adenoma and carcinoma patients with nuclear β -catenin expression (IHC scores) in their respective lesions.

	Patients	Sex	Age	Diagnosis	Duke's	Size of	Plasma	IHC of
				•	stage	lesion	β-catenin	β-catenin
	1	F	65	granulation tissue	N.A.	N.A.	•	+
	2	F	68	granulation tissue	N.A.	N.A.	_	-
20	3	F	59	dilated lymphatic space	N.A.	N.A.	-	-
	4	F	68	adenoma, moderate dys	N.A.	N.A.	+	+
	5	F	75	adenoma, mild dys	N.A.	N.A.	+	++
	6	M	82	adenoma, mild dys	N.A.	N.A.	+	+
	7	F	61	adenoma, moderate dys	N.A.	5 mm	+	+
25	8	M	68	adenoma, moderate dys	N.A.	4 mm	+	+
	9	F	77	adenoma, moderate dys	N.A.	N.A.	+	}
	10	M	72	adenoma, moderate dys	N.A.	10 mm	-	1-1
	11	M	51	adenoma, mild dys	N.A.	N.A.	+	+
	12	F	81	adenoma, moderate dys	N.A.	N.A.	+	+
30	13	M	67	adenoma, moderate dys	N.A.	72 mm ³	+	++
	14	M	75	adenoma, moderate dys	N.A.	672 mm ³	-	++
	15	M	70	adenoma, mild dys	N.A.	N.A.	+	+
	16	M	78	adenoma, severe dys	N.A.	1500 mm ³	-	+++
	17	F	73	adenoma, severe dys	N.A.	1200 mm ³	+	+++
35	18	M	59	adenocarcinoma	В	91 cm ³	+	+
	19	F	56	adenocarcinoma	C	90 cm ³	+	+
	20	F	67	adenocarcinoma	С	108 cm ³	+	+
	21	F	75	adenocarcinoma	С	$100 \mathrm{cm}^3$	+	++++
	22	F	92	adenocarcinoma	N.A.	N.A.	+	++++
40	23	F	79	adenocarcinoma	N.A.	N.A.	+	++
	24	F	76	adenocarcinoma	В	88 cm ³	+	++
	25	M	82	adenocarcinoma	D	115 cm ³	+	+
	26	F	77	adenocarcinoma	В	346 cm ³	+	+
	27	F	73	adenocarcinoma	Α	$21 \mathrm{cm}^3$	+	++++
45	28	F	82	adenocarcinoma	N.D.	N.D.	+	++++
	29	F	80	adenocarcinoma	В	130 cm ³	+	++
	30	M	77	adenocarcinoma	В	155 cm ³	+	+++
		M	62	adenocarcinoma	В	167 cm ³	+	+-+
	32	F	85	adenocarcinoma	В	143 cm^3	+	+++
50	3	1	NT A	not applied MID at date				

dys: dysplasia; N.A.: not applied; N.D.: not determined.

Table 3: Primers used in the PCR reactions.

	Primer	Nucleotide sequence (5' to 3')	Design P	roduct size
5	1 sen	se: TCAATGGGTCATATCACAGAT isense: CTGCATTCTGACTTTCAGTAA	In intron 2 and 3 of β-Catenin gene	359 bp
	2 sen	se: CCTCTGCGGTGCCAAGCCTC isense: TGTGGGCAAACTTGTGGTAGCA	Within exon 11 of RET gene	e 156 bp

Table 4. Patients record

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Patient 1 2 3	Sex M F	Age 23	Diagnosis Du adenoma, severe dys	ike's stage	Size of lesion
2 3			adenoma severe due		
3	F		auchoma, severe dys	N.A.	75mm ³
		48	adenoma, moderate dys	N.A.	N.A.
	M	67	adenoma, moderate dys	N.A.	168mm ³
4	M	67	adenoma, severe dys	N.A.	$80 \mathrm{mm}^3$
5	M	76	adenoma, severe dys	N.A.	63mm ³
6	F	62	adenoma, mild dys	N.A.	N.A.
7	M	85	adenoma, severe dys	N.A.	153mm ³
8	F	81	adenoma, moderate dys	N.A.	96mm ³
9	F	58	adenoma, moderate dys	N.A.	48mm ³
10	F	68	adenoma, moderate dys	N.A.	528mm ³
11	M	62	adenocarcinoma	В	182cm^3
12	M	67	adenocarcinoma	В	72cm ³
13	M	83	adenocarcinoma	В	43cm ³
14	M	45	adenocarcinoma	C	$67 \mathrm{cm}^3$
15	M	52	adenocarcinoma	С	41cm ³
16	F	71	adenocarcinoma	C	64cm ³
17	M	80	adenocarcinoma	C .	47cm ³
18	M	61	adenocarcinoma	N.D.	N.A.
19	F	70	adenocarcinoma	Α	13cm ³
20	M	69	adenocarcinoma	В	$120 \mathrm{cm}^3$
21	M	61	adenocarcinoma	С	384cm ³
22	F	72	adenocarcinoma	Α	9cm³
23	M	76	adenocarcinoma	N.D.	N.A.
24	M	76	adenocarcinoma	С	88cm ³
25	M	70	adenocarcinoma	В	$23 cm^3$
	7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24	6 F 7 M 8 F 9 F 10 F 11 M 12 M 13 M 14 M 15 M 16 F 17 M 18 M 19 F 20 M 21 M 22 F 23 M 24 M	6 F 62 7 M 85 8 F 81 9 F 58 10 F 68 11 M 62 12 M 67 13 M 83 14 M 45 15 M 52 16 F 71 17 M 80 18 M 61 19 F 70 20 M 69 21 M 61 22 F 72 23 M 76 24 M 76	6 F 62 adenoma, mild dys 7 M 85 adenoma, severe dys 8 F 81 adenoma, moderate dys 9 F 58 adenoma, moderate dys 10 F 68 adenoma, moderate dys 11 M 62 adenocarcinoma 12 M 67 adenocarcinoma 13 M 83 adenocarcinoma 14 M 45 adenocarcinoma 15 M 52 adenocarcinoma 16 F 71 adenocarcinoma 17 M 80 adenocarcinoma 18 M 61 adenocarcinoma 19 F 70 adenocarcinoma 20 M 69 adenocarcinoma 21 M 61 adenocarcinoma 22 F 72 adenocarcinoma 23 M 76 adenocarcinoma 24 M 76 adenocarcinoma	6 F 62 adenoma, mild dys N.A. 7 M 85 adenoma, severe dys N.A. 8 F 81 adenoma, moderate dys N.A. 9 F 58 adenoma, moderate dys N.A. 10 F 68 adenoma, moderate dys N.A. 11 M 62 adenocarcinoma B 12 M 67 adenocarcinoma B 13 M 83 adenocarcinoma B 14 M 45 adenocarcinoma C 15 M 52 adenocarcinoma C 16 F 71 adenocarcinoma C 17 M 80 adenocarcinoma C 18 M 61 adenocarcinoma C 19 F 70 adenocarcinoma A 20 M 69 adenocarcinoma B 21 M 61 adenocarcinoma C 22 F 72 adenocarcinoma C 23 M 76 adenocarcinoma N.D. 24 M 76 adenocarcinoma C

dys: dysplasia; N.A.: not applied; N.D.: not determined

While various embodiments are disclosed herein, it should be understood that they have been presented by way of example only, and not limitation. Thus, the breadth and scope of the invention(s) should not be limited by any of the above-described exemplary embodiments, but should be defined only in accordance with the following claims and their equivalents. Moreover, the above advantages and features are affected in described embodiments, but shall not limit the application of the claims to processes and structures accomplishing any or all of the above advantages.

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Furthermore, teachings from the following references are incorporated herein by reference for all purposes:

- Anker, P., Lefort, F., Vasioukhin, V., Lyautey, J., Lederrey, C., Chen, X.Q., Stroun, M., Mulcahy, H.E. and Farthing, M.J. K-ras mutations are found in DNA extracted from the plasma of patients with colorectal cancer. Gastroenterology 112: 1114-1120, 1997.
- Chen, X. Q., Bonnefoi, H., Pelte, M-F., Lyautey, J., Lederrey, C., Movarekhi, S., Schaeffer, P., Mulcahy, H. E., Meyer, P., Stroun, M. and Anker, P. Telomerase RNA as a detection marker in the serum of breast cancer patients. Clinical Cancer Research 6: 3823-3826, 2000.
- Hibi, K., Robinson, C.R., Booker, S., Wu, L., Hamilton, S.R., Sidransky, D. and Jen, J. Molecular detection of genetic alterations in the serum of colorectal cancer patients. 58: 1405-1407, 1998.
- Kopreski, M. S., Benko, F. A., Kwak, L. W. and Gocke, C. D. Detection of tumor suppressor messenger RNA in the serum of patients with malignant melanoma. Clinical Cancer Research 5: 1961-1965, 1999.
 - Kopreski, M.S., Benko, F.A. and Gocke, C.D. Circulating RNA as a tumor marker: detection of 5T4 mRNA in breast and lung cancer patient serum. Ann. N.Y. Acad. Sci. 945: 172-178, 2001.
- 20 Lo, K. W., Lo, Y. M. D., Leung, S. F., Tsang, Y. S., Chan, L. Y. S., Johnson, P. J., Hjelm, N. M., Lee, J. C. K. and Huang, D. P. Analysis of cell-free Epstein-Barr virus-associated RNA in the plasma of patients with nasopharyngeal carcinoma. Clinical Chemistry 45: 1292-1294, 1999.
- Matias-Guiu, X. RET protooncogene analysis in the diagnosis of medullary thyroid carcinoma and multiple endocrine neoplasia type II. Advances in Anatomic Pathology 5: 196-201, 1998.
 - Morin, P.J. β-catenin signaling and cancer. Bioessays, 21: 1021-1030, 1999. Remmele, W., Schicketanz, K.H. Immunohistochemical determination of estrogen and progesterone receptor content in human breast cancer. Computerassisted image analysis (QIC score) vs subjective grading IRS. Pathol Res Pract 189: 862-866, 1993.
 - Sozzi, G., Musso, K., Ratcliffe, C., Goldstraw, P., Pierotti, M.A. and Pastorino, U. Detection of microsatellite alterations in plasma DNA of non-small cell lung

cancer patients: a prospect for early diagnosis. Clin. Cancer Res. 5: 2689-2692, 1999.

von Knobloch, R., Hegele, A., Brandt, H., Olbert, P., Heidenreich, A. and Hofman, R. Serum DNA and urine DNA alterations of urinary transitional cell bladder carcinoma detected by fluorescent microsatellite analysis. Int.J. Cancer 94: 67-72, 2001.

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Willert, K. and Nusse, R. β-catenin: a key mediator of Wnt signaling. Curr. Opin, Genet. Dev. 8: 95-102, 1998.

Wong, S.C., Chan, K.C., Lee, K.C., Hsiao, W.L. Differential expressions of p16/p21/p27 and cyclin D1/D3, and their relationships to cell proliferation, apoptosis and tumor progression in invasive breast ductal carcinoma. J Pathol 194: 35-42, 2001.

Additionally, the section headings herein are provided for consistency with the suggestions under 37 CFR 1.77 or otherwise to provide organizational cues. These headings shall not limit or characterize the invention(s) set out in any claims that may issue from this disclosure. Specifically and by way of example, although the headings refer to a "Technical Field of the Invention," the claims should not be limited by the language chosen under this heading to describe the so-called field of the invention. Further, a description of a technology in the "Background of the Invention" is not to be construed as an admission that technology is prior art to any invention(s) in this disclosure. Neither is the "Brief Summary of the Invention" to be considered as a characterization of the invention(s) set forth in the claims set forth herein. Furthermore, the reference in these headings, or elsewhere in this disclosure, to "invention" in the singular should not be used to argue that there is only a single point of novelty claimed in this disclosure. Multiple inventions may be set forth according to the limitations of the multiple claims associated with this disclosure, and the claims, and their equivalents, accordingly define the invention(s) that are protected thereby. In all instances, the scope of the claims shall be considered on their own merits in light of the specification, but should not be constrained by the headings set forth herein.

WHAT IS CLAIMED IS:

- 1. A method for detecting non-clinically diagnosed cancer in a patient, the method comprising:
- extracting blood serum or plasma from the patient;

 detecting beta-catenin RNA in the blood serum or plasma; and

 determining the presence of the cancer based on the detected beta-catenin

 RNA.
- 2. A method according to claim 1, wherein determining the presence of the cancer comprises determining the presence of colorectal cancer based on the detected beta-catenin RNA.
- A method according to claim 2, wherein determining the presence of
 colorectal cancer comprises detecting pre-neoplastic colorectal polyps based on the detected beta-catenin RNA.
 - 4. A method according to claim 1, wherein the RNA is derived from one of the group consisting of:
- 20 gene-encoded beta-catenin,
 gene-encoded alpha-catenin,
 gene-encoded E-catherin, and
 other gene-encoded beta-catenin associated proteins.
- 25 5. A method according to claim 1, wherein the patient is a human or animal.
 - 6. A method for detecting non-clinically diagnosed cancer in a patient, the method comprising:

extracting blood serum or plasma from the patient;

detecting beta-catenin DNA in the blood serum or plasma; and
determining the presence of the cancer based on the detected beta-catenin DNA.

7. A method according to claim 6, wherein determining the presence of the cancer comprises determining the presence of colorectal cancer based on the detected beta-catenin DNA.

- 5 8. A method according to claim 7, wherein determining the presence of colorectal cancer comprises detecting pre-neoplastic colorectal polyps based on the detected beta-catenin DNA.
- 9. A method according to claim 6, wherein the DNA is derived from one of the10 group consisting of:

gene-encoded beta-catenin,
gene-encoded alpha-catenin,
gene-encoded E-catherin, and
other gene-encoded beta-catenin associated proteins.

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- 10. A method according to claim 6, wherein the patient is a human or animal.
- 11. A method for detecting non-clinically diagnosed cancer in a patient, the method comprising:
- 20 extracting blood serum or plasma from the patient;
 detecting beta-catenin-associated gene RNA in the blood serum or plasma; and
 determining the presence of the cancer based on the detected beta-cateninassociated gene RNA.
- 25 12. A method according to claim 11, wherein determining the presence of the cancer comprises determining the presence of colorectal cancer based on the detected beta-catenin-associated gene RNA.
- 13. A method according to claim 12, wherein determining the presence of
 30 colorectal cancer comprises detecting pre-neoplastic colorectal polyps based on the detected beta-catenin-associated gene RNA.
 - 14. A method according to claim 11, wherein the RNA is derived from one of the group consisting of:

gene-encoded beta-catenin,
gene-encoded alpha-catenin,
gene-encoded E-catherin, and
other gene-encoded beta-catenin associated proteins.

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- 15. A method according to claim 11, wherein the patient is a human or animal.
- 16. A method for detecting non-clinically diagnosed cancer in a patient, the method comprising:

extracting blood serum or plasma from the patient;

detecting beta-catenin-associated gene DNA in the blood serum or plasma; and

determining the presence of the cancer based on the detected beta-cateninassociated gene DNA.

17. A method according to claim 16, wherein determining the presence of the cancer comprises determining the presence of colorectal cancer based on the detected beta-catenin-associated gene DNA.

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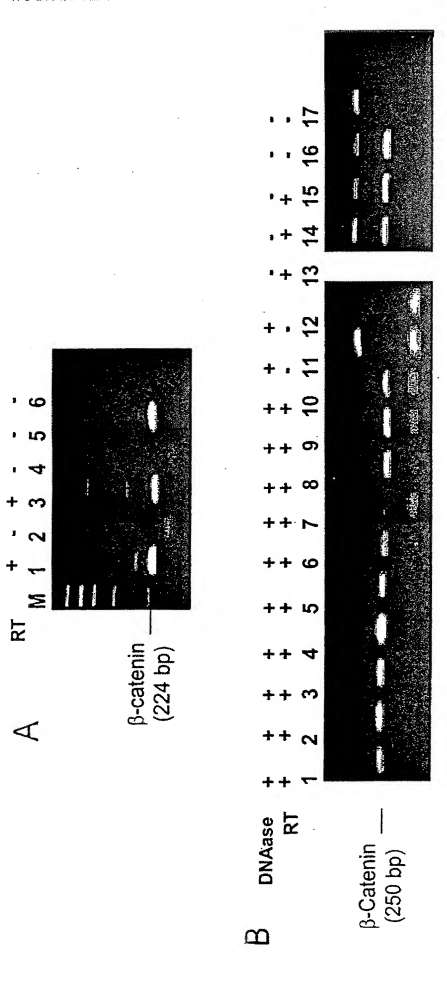
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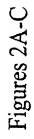
- 18. A method according to claim 17, wherein determining the presence of colorectal cancer comprises detecting pre-neoplastic colorectal polyps based on the detected beta-catenin-associated gene DNA.
- 25 19. A method according to claim 16, wherein the DNA is derived from one of the group consisting of:

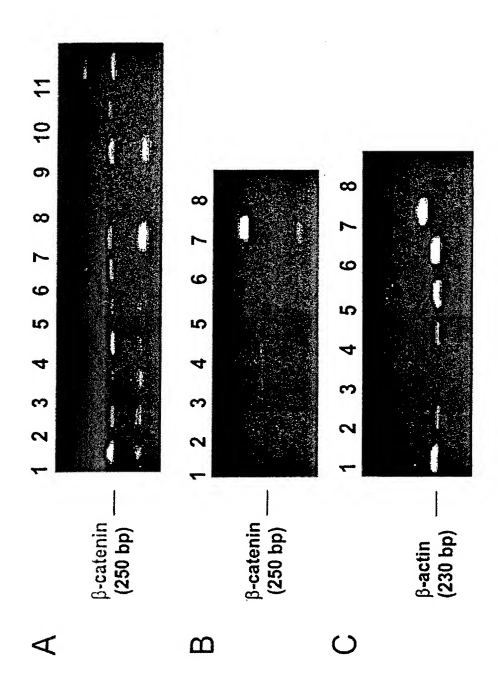
gene-encoded beta-catenin, gene-encoded alpha-catenin, gene-encoded E-catherin, and

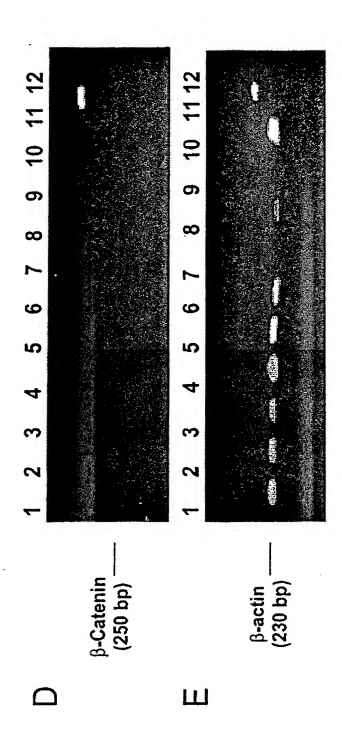
- other gene-encoded beta-catenin associated proteins.
 - 20. A method according to claim 16, wherein the patient is a human or animal.

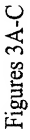


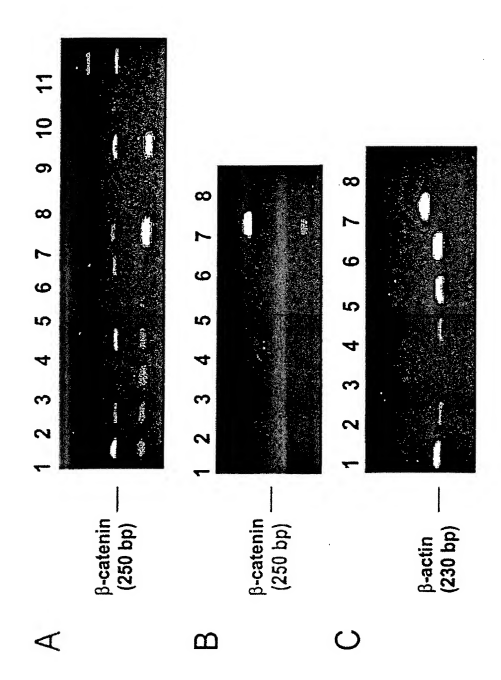


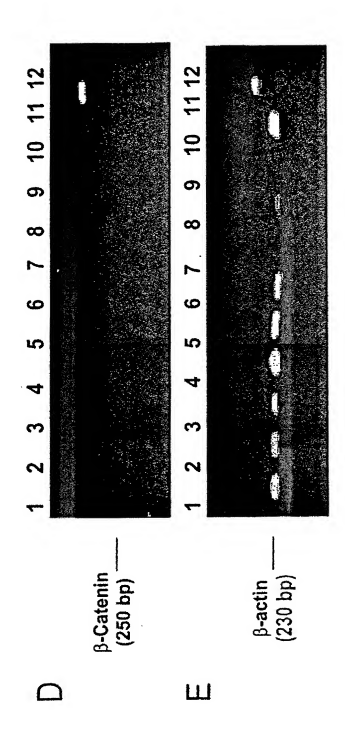












AMENDED CLAIMS

[received by the International Bureau on 08 April 2004 (08.04.04) original claims 1-20 have been amended, 21-29 have been added.]

WHAT IS CLAIMED IS:

- A method for detecting cancer in a patient, comprising:
 extracting blood serum or plasma from the patient;
 detecting the presence or absence of beta-catenin RNA in the blood serum of plasma;
 and
 determining the presence of the cancer based on the detected presence of beta-catenin
 RNA.
- 2. The method according to claim 1, whereby the cancer is colorectal cancer.
- 3. The method according to claim 2, whereby determining the presence of colorectal cancer comprises detecting pre-neoplastic colorectal polyps based on the detected presence of beta-catenin RNA.
- 4. The method according to claim 1, whereby the RNA is derived from one of the group consisting of:

gene-encoded beta-catenin,
gene-encoded alpha-catenin,
gene-encoded E-catherin, and
other gene-encoded beta-catenin associated proteins.

- 5. The method according to claim 1, whereby the patient is a human or animal.
- A method for detecting cancer in a patient, comprising:
 extracting blood serum or plasma from the patient;
 detecting the presence or absence of beta-catenin DNA in the blood serum or plasma;
 and

determining the presence of the cancer based on the detected presence of beta-catenin DNA.

- The method according to claim 6, whereby the cancer is colorectal cancer.
- 8. The method according to claim 7, whereby determining the presence of colorectal cancer comprises detecting pre-neoplastic colorectal polyps based on the detected presence of beta-catenin DNA.
- 9. The method according to claim 6, whereby the DNA is derived from one of the group consisting of:

gene-encoded beta-catenin,

gene-encoded alpha-catenin,
gene-encoded E-catherin, and
other gene-encoded beta-catenin associated proteins.

- 10. The method according to claim 6, whereby the patient is a human or animal.
- A method for detecting cancer in a patient, comprising:
 extracting blood serum or plasma from the patient;

detecting the presence or absence of beta-catenin-associated gene RNA in the blood serum or plasma; and

determining the presence of the cancer based on the detected presence of beta-catenin associated gene RNA.

- 12. The method according to claim 11, whereby the cancer is colorectal cancer.
- 13. The method according to claim 12, whereby determining the presence of colorectal cancer comprises detecting pre-neoplastic colorectal polyps based on the detected beta-catenin-associated gene RNA.
- 14. The method according to claim 11, whereby the RNA is derived from one of the group consisting of:

gene-encoded beta-catenin,
gene-encoded alpha-catenin,
gene-encoded E-catherin, and
other gene-encoded beta-catenin associated proteins.

- 15. The method according to claim 11, whereby the patient is a human or animal.
- 16. A method for detecting cancer in a patient, comprising: extracting blood serum or plasma from the patient;

detecting the presence or absence of beta-catenin-associated gene DNA in the blood serum or plasma; and

determining the presence of the cancer based on the detected presence of beta-cateninassociated gene DNA.

17. The method according to claim 16, whereby the cancer is colorectal cancer.

18. The method according to claim 17, whereby determining the presence of colorectal cancer comprises detecting pre-neoplastic colorectal polyps based on the presence of detected beta-catenin-associated gene DNA.

19. The method according to claim 16, whereby the DNA is derived from one of the group consisting of:

gene-encoded beta-catenin,
gene-encoded alpha-catenin,
gene-encoded E-catherin, and
other gene-encoded beta-catenin associated proteins.

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- 20. The method according to claim 16, whereby the patient is a human or animal.
- 21. The method according to claims 2, 7, 12, or 16, whereby the colorectal cancer is colorectal carcinoma or colorectal adenoma.
- 22. A method of determining the presence of carcinoma, the presence of adenoma, or the absence of carcinoma and adenoma in a patient, comprising:

extracting blood serum or plasma from a patient,

measuring the relative amount of beta-catenin DNA or RNA in the blood serum or plasma of the patient and the relative amount of beta-catenin DNA or RNA in the blood serum or plasma of a control person known not to have carcinoma or adenoma;

determining a ratio of the amount of beta-catenin DNA or RNA detected in the blood serum or plasma of the patient to the amount of beta-catenin DNA or RNA detected in the blood serum or plasma of a control person known not to have carcinoma or adenoma, whereby the ratio of approximately 30-80 indicates the presence of adenoma, the ratio of approximately above 500 indicates the presence of carcinoma, and the ratio of approximately 1 indicates the absence of carcinoma and adenoma.

- 23. The method according to claim 22, whereby the carcinoma is colorectal carcinoma.
- 24. The method according to claim 22, whereby the adenoma is colorectal adenoma.
- 25. The method according to claim 22, whereby the DNA or RNA is derived from one of the group consisting of:

gene-encoded beta-catenin, gene-encoded alpha-catenin,

to a second of the

gene-encoded E-catherin, and other gene-encoded beta-catenin associated proteins.

- 26. The method according to claim 22, whereby the ratio of 30 indicates the presence of adenoma.
- 27. The method according to claim 22, whereby the ratio of 598 indicates the presence of carcinoma.
- 28. The method according to claim 22, whereby the relative amount of beta-catenin DNA or RNA in the blood serum or plasma of the patient and the relative amount of beta-catenin DNA or RNA in the blood serum or plasma of a control person known not to have carcinoma or adenoma is measured using real time reverse transcription-polymerase chain reactions.
- 29. The method according to claims 1, 6, 11, or 16, whereby the detecting step is accomplished using reverse transcription-polymerase chain reactions (RT-PCR).

Exhibit C

Leung, Monica

寄件者: 寄件日期: 收件者:

Cesar Wong [cesar01@netvigator.com]

Friday, August 26, 2005 9:45 PM

主旨:

Cheung, Peggy
Read: HKUST - US Patent Application "Plasma or Serum Marker and Process for

Detection of Cancer" (Our Ref: 32144183-000004)

附件:

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This is a receipt for the mail you sent to <cesar01@netvigator.com> at 8/26/2005 5:49 PM

This receipt verifies that the message has been displayed on the recipient's computer at 8/26/2005 9:44 PM

DELIVERY / PICK UP SERVICES REQUEST FORM

To: OFFICE SERVICES	Date: 30 August 2005 Time. 2:30 pm
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Client/Matter No.: 32144183-000004 Conve	1g No.:
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Address: Flat C, Floor 9, King's Court, 14-16 Vil	King's Court, 14-16 Village Road, Happy Valley, Hong Kong
医要抗 Delive	Deliver By Date/Time: 30 August 2005 at 9:00pm
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Exhibit E2

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Customer Service Hotline: (852)2710-8111

Detail Report for Shipment: 1048694091

Status	Time	Date
Shipment received from sender	12:35 p.m.	2/9/2005
Delivering shipment	02:02 p.m.	2/9/2005

Go to Top

Exhibit E-3

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